

This Page Is Inserted by IFW Operations
and is not a part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images may include (but are not limited to):

- BLACK BORDERS
- TEXT CUT OFF AT TOP, BOTTOM OR SIDES
- FADED TEXT
- ILLEGIBLE TEXT
- SKEWED/SLANTED IMAGES
- COLORED PHOTOS
- BLACK OR VERY BLACK AND WHITE DARK PHOTOS
- GRAY SCALE DOCUMENTS

IMAGES ARE BEST AVAILABLE COPY.

**As rescanning documents *will not* correct images,
please do not report the images to the
Image Problem Mailbox.**

THIS PAGE BLANK (USPTO)

DNG-1775
PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION
International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5 : A61K 31/557 // (A61K 31/557 A62K 31:415)		A1	(11) International Publication Number: WO 94/08585 (43) International Publication Date: 28 April 1994 (28.04.94)
(21) International Application Number: PCT/US93/09742 (22) International Filing Date: 12 October 1993 (12.10.93)		(81) Designated States: AU, CA, JP, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE). Published <i>With international search report.</i>	
(30) Priority data: 07/960,065 13 October 1992 (13.10.92) US			
(71) Applicant: ALCON LABORATORIES, INC. [US/US]; 6201 South Freeway, Fort Worth, TX 76134 (US).			
(72) Inventors: DESANTIS, Louis, Jr. ; 2316 Winton Terrace West, Fort Worth, TX 76109 (US). SALLEE, Verney, L. ; 304 Diamond Lane Road, Burleson, TX 76028 (US).			
(74) Agents: CHENG, Julie et al.; Alcon Laboratories, Inc., 6201 South Freeway, Fort Worth, TX 76134 (US).			
(54) Title: COMBINATIONS OF PROSTAGLANDINS AND CLONIDINE DERIVATIVES FOR THE TREATMENT OF GLAUCOMA			
(57) Abstract Combinations of at least one clonidine derivative and at least one prostaglandin are used to treat glaucoma and ocular hypertension without some of the side effects typically associated with topical administration of prostaglandins.			

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AT	Austria	FR	France	MR	Mauritania
AU	Australia	GA	Gabon	MW	Malawi
BB	Barbados	GB	United Kingdom	NE	Niger
BE	Belgium	GN	Guinea	NL	Netherlands
BF	Burkina Faso	GR	Greece	NO	Norway
BG	Bulgaria	HU	Hungary	NZ	New Zealand
BJ	Benin	IE	Ireland	PL	Poland
BR	Brazil	IT	Italy	PT	Portugal
BY	Belarus	JP	Japan	RO	Romania
CA	Canada	KP	Democratic People's Republic of Korea	RU	Russian Federation
CF	Central African Republic	KR	Republic of Korea	SD	Sudan
CC	Congo	KZ	Kazakhstan	SE	Sweden
CH	Switzerland	LI	Liechtenstein	SI	Slovenia
CI	Côte d'Ivoire	LK	Sri Lanka	SK	Slovak Republic
CM	Cameroon	LU	Luxembourg	SN	Senegal
CN	China	LV	Latvia	TD	Chad
CS	Czechoslovakia	MC	Monaco	TG	Togo
CZ	Czech Republic	MG	Madagascar	UA	Ukraine
DE	Germany	ML	Mali	US	United States of America
DK	Denmark	MN	Mongolia	UZ	Uzbekistan
ES	Spain			VN	Viet Nam
FI	Finland				

COMBINATIONS OF PROSTAGLANDINS AND CLONIDINE DERIVATIVES FOR THE TREATMENT OF GLAUCOMA

BACKGROUND OF THE INVENTION

1. Field of the Invention

5 The present invention relates generally to the field of ophthalmology. In particular, the invention relates to the treatment of glaucoma and ocular hypertension using a combination of at least one clonidine derivative (e.g., para-amino clonidine) and at least one prostaglandin.

2. Discussion of Related Art

10 Although the underlying causes of glaucoma are not understood, its symptoms often include elevated intraocular pressure, which may be caused either by over-production of aqueous humor or by inadequate outflow of aqueous humor. If left untreated, or if inadequately treated, glaucoma can lead to blindness or significant loss of vision. There is therefore a continuing need for therapies which control the elevated intraocular pressure associated with glaucoma.

15 There are currently a number of drugs utilized in the treatment of glaucoma, including: miotics (e.g., pilocarpine, carbachol and acetylcholinesterase inhibitors); sympathomimetics (e.g., epinephrine and dipivalylepinephrine); alpha-2 agonists (e.g., para-amino clonidine); beta-blockers (e.g., betaxolol, levobunolol and timolol); and carbonic anhydrase inhibitors (e.g., acetazolamide, methazolamide and ethoxzolamide). Miotics and sympathomimetics are believed to lower IOP by increasing the outflow of aqueous humor through the trabecular meshwork, while beta-blockers, alpha-2 agonists and carbonic anhydrase inhibitors are believed to lower IOP by decreasing the formation of aqueous humor.

In addition, although they have not yet been approved for anti-glaucoma therapy, certain classes of prostaglandins and prostaglandin analogues (hereinafter collectively referred to as "prostaglandins") have been shown in various animal models and in some clinical studies to reduce intraocular pressure (IOP) to a greater extent than most currently used therapeutic agents. See, for example: US 4,097,489 (Bundy), US 4,599,353 (Bito), US 4,994,274 (Chan et al.) and EP 289 349 (Ueno et al.). In contrast to the case with miotics, prostaglandins are believed to lower IOP by increasing the outflow of aqueous humor via the uveo-scleral route. In addition, prostaglandins may possibly have other effects in the eye, such as enhancing vascular support of ocular tissues; however, there is no understanding 10 of that mechanism at this time.

All six types of therapeutic agents have potentially serious side effects: miotics such as pilocarpine can cause blurring of vision and other, visual side effects, which may lead either to decreased patient compliance or to termination of 15 therapy; carbonic anhydrase inhibitors can also cause serious side effects which affect patient compliance and/or necessitate the withdrawal of treatment; at least one beta-blocker, timolol, has increasingly become associated with serious pulmonary side effects attributable to its effect on beta-2 receptors in pulmonary tissue; and prostaglandins often produce hyperemia and edema of the conjunctiva, resulting in redness and hyperesthesia of the eye, which may affect patient 20 compliance. In addition to these side effects, a therapy regimen which includes the use of two or more pharmaceutical compositions containing drugs selected from two or more of the above-cited classes requires the patient to apply the compositions to the affected eye(s) in separate, spaced dosages, several times per day. Patient compliance with such complicated dosage regimens can be very poor, particularly in elderly patients. Since the majority of glaucoma patients are elderly, 25 this patient compliance problem is significant.

In light of the foregoing circumstances, it is clear that a need exists for new, more potent anti-glaucoma compositions which avoid or reduce the above-cited

side effects, while increasing patient compliance. The present invention is directed to such compositions.

SUMMARY OF THE INVENTION

It has unexpectedly been found that administration of one or more prostaglandins in combination with one or more clonidine derivatives controls or lowers intraocular pressure (IOP) without the accompanying inflammatory response (including hyperemia) typically found with prostaglandins. The present invention therefore provides compositions and methods useful for the treatment of glaucoma and ocular hypertension. The compositions contain a combination of at least one clonidine derivative and at least one prostaglandin which is effective in reducing or controlling IOP, and which has a reduction or elimination of the side effects normally associated with topical application of prostaglandins.

In a preferred formulatory embodiment of the compositions of the present invention, the above combinations may further include an anionic mucomimetic polymer, a gelling polysaccharide, a finely divided drug carrier substrate (defined below), or a combination of these components. These additional components provide compositions which are comfortable and have sustained release.

DETAILED DESCRIPTION OF THE INVENTION

The present invention utilizes combinations of at least one clonidine derivative and at least one prostaglandin to treat glaucoma and ocular hypertension.

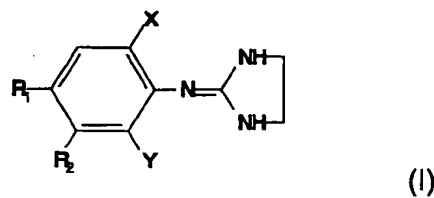
Clonidine is a known hypotensive compound, and is described, for example, in US 3,202,660 (Zeile et al.); the contents of this patent relating to the structure, preparation and physical properties of this compound are incorporated herein by

-4-

reference. It is also known that certain clonidine derivatives are effective in lowering intraocular pressure when applied topically to the eye; this discovery is described in US 4,461,904 (York, Jr.), the entire contents of which are incorporated herein by reference. The clonidine derivatives described in this patent are 2-(tri-substituted phenylimino)-imidazoline compounds, which are also known as 2-(tri-substituted anilino)-1,3-diazacyclopentene-(2) compounds. Reference is made to this patent for further details concerning the structure, preparation and physical properties of these clonidine derivatives. Related developments are described in US 4,517,199 (York, Jr.), US 4,587,257 (DeSantis et al.) and US 4,515,800 (Cavero et al.); which are all incorporated herein by reference to the extent that they disclose, generically and specifically, the subject clonidine-like compounds.

A comprehensive discussion of the properties of clonidine and clonidine-like compounds is presented in a publication by Timmermans et al. entitled: "Structure-Activity Relationships in Clonidine-Like Imidazolidines and Related Compounds" (Gustav Fischer Verlag, New York: 1980, page 1-97). The entire contents of that publication are incorporated herein by reference. As indicated by Timmermans et al., the molecular structure of clonidine consists of three parts: an aromatic (i.e., aryl) portion, a bridge, and an imidazolidine moiety. Timmermans et al. disclose many compounds which have been produced by modifying one or two of these three parts, but which retain one of the three parts intact. For purposes of the present specification, all such compounds are defined as being "clonidine derivatives."

A preferred group of clonidine derivatives are those of formula:



25

wherein: R₁ and R₂ are selected from H, OH, NHR' and O-C(=O)-CH₂-R',

-5-

with R' being selected from H and C₁-C₄ alkyl, provided that one of R₁ and R₂ is hydrogen; and X and Y are selected from Br, Cl, CH₃ and CH₂CH₃.

Specific examples of Compounds from this group are set forth in Table 1, below.

TABLE 1

5

Compound	R ₁	R ₂	X	Y
1	NHCH ₃	H	CH ₃	CH ₃
2	NHCH ₃	H	CH ₂ CH ₃	CH ₂ CH ₃
3	NHCH ₃	H	Cl	Cl
4	NH ₂	H	Br	Br

A group of especially preferred clonidine derivatives of formula (I) are those in which R₁ and R₂ are selected from H and NH₂, provided that one of R₁ and R₂ is H, and X and Y are selected from Cl, CH₃, and CH₂CH₃. Specific examples of compounds from this group are set forth in Table 2, below.

TABLE 2

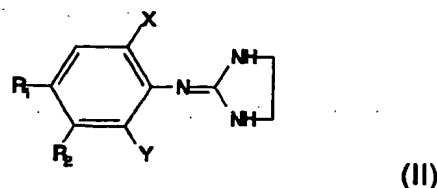
15

Compound	R ₁	R ₂	X	Y
5	H	NH ₂	CH ₃	CH ₃
6	NH ₂	H	CH ₂ CH ₃	CH ₂ CH ₃
7	H	NH ₂	Cl	Cl
8	NH ₂	H	CH ₂ CH ₃	Cl
9	NH ₂	H	CH ₃	Cl
10	NH ₂	H	CH ₂ CH ₃	CH ₃
11	NH ₂	H	CH ₃	CH ₃
12	H	NH ₂	CH ₂ CH ₃	CH ₂ CH ₃
13	NH ₂	H	Cl	Cl

25

Of these specific examples, Compound 13 para-amino clonidine (also known as apraclonidine), has been found to be particularly well-suited for use in the present invention.

5 A second preferred group of clonidine derivatives useful in the present invention are those of formula:



wherein: X and Y are selected from Br, Cl, CH₃ and CH₂CH₃, with the provision that at least one of X and Y is alkyl. Compounds of this type are described, for example in US 3,468,887 (Stahle et al.), and J. Med. Chem., 19: 1049-54 (1976).

10 The contents of this patent and article relating to the structure, preparation and physical properties of these compounds are incorporated herein by reference. Specific examples of compounds from this group are set forth in Table 3, below.

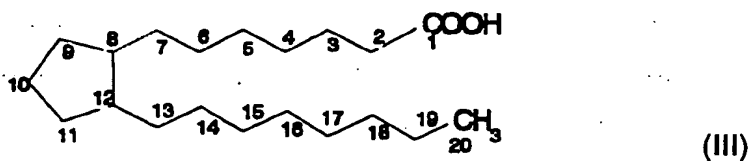
TABLE 3

Compound	X	Y
14	CH ₂ CH ₃	CH ₂ CH ₃
15	CH ₂ CH ₃	CH ₃
16	Cl	CH ₂ CH ₃

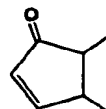
In addition to the 2-(arylimino) imidazolidines identified above, other groups or classes of alpha-2 agonists which may be utilized in the present invention include 2-(arylimino) oxazolidines; 2-(aryl)methylene) imidazolidines; 2-(arylimino) pyrrolidines; arylalkylaminoguanidines, such as aryl-imidazoquinazolines and phenyl-acetyguanidines; and 2-(phenylimino) diazocyclopentenes. All of these groups of drugs may be referred to as being clonidine derivatives or "clonidine-like" drugs.

-7-

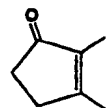
The terms "prostaglandin" and "PG" are generally used to describe a class of compounds which are analogues and derivatives of prostanoic acid (III):



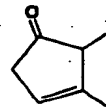
PG's may be further classified, for example, according to their 5-membered ring structure, using a letter designation:



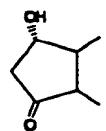
Prostaglandins of the A series (PGA's):



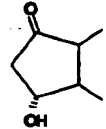
Prostaglandins of the B series (PGB's):



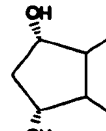
Prostaglandins of the C series (PGC's):



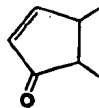
Prostaglandins of the D series (PGD's):



Prostaglandins of the E series (PGE's):



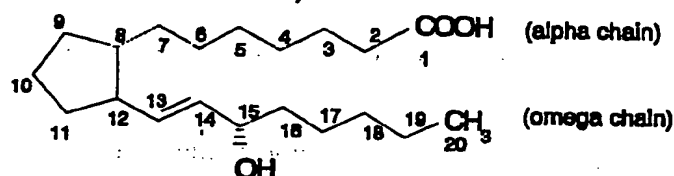
Prostaglandins of the F series (PGF's):



Prostaglandins of the J series (PGJ's):

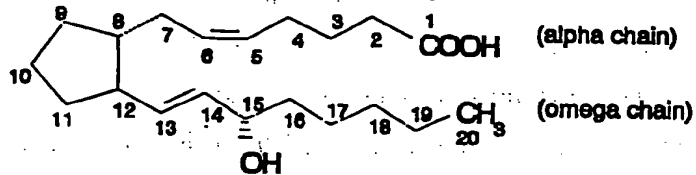
PG's may be further classified based on the number of unsaturated bonds on the side chain:

PG's (13,14- unsaturated):

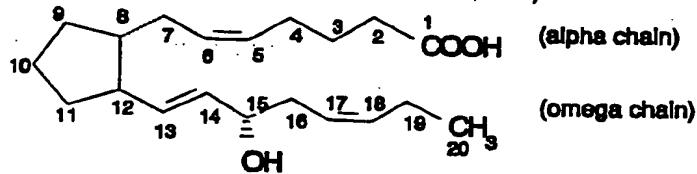


5

PG₂'s (13,14- and 5,6- unsaturated):



PG₃'s (13,14- 5,6- and 17,18- unsaturated):



10

An historical review of the ocular effects of prostaglandins and other eicosanoids can be found in Bito, L. and J. Stjernschantz, The Ocular Effect of Prostaglandins and Other Eicosanoids, Alan R. Liss, Inc., New York:1989, 1-13.

The prostaglandins which may be utilized in the present invention include all pharmaceutically acceptable prostaglandins, their derivatives and analogues, and their pharmaceutically acceptable ester and salts (hereinafter collectively referred to as "prostaglandins" or "PG's"), which are capable of reducing intraocular pressure when applied topically to the eye. Such prostaglandins include the natural compounds: PGE₁, PGE₂, PGE₃, PGF_{1 α} , PGF_{2 α} , PGF_{3 α} , PGD₂ and PGI₂

15.

(prostacyclin), as well as analogues and derivatives of these compounds which have similar biological activities of either greater or lesser potencies. Analogues of the natural prostaglandins include but are not limited to: alkyl substitutions (e.g., 15-methyl or 16,16-dimethyl), which confer enhanced or sustained potency by reducing biological metabolism or alter selectivity of action; saturation (e.g., 13,14-dihydro) or unsaturation (e.g., 2,3-didehydro, 13,14-didehydro), which confer sustained potency by reducing biological metabolism or alter selectivity of action; deletions or replacements (e.g., 11-deoxy, 9-deoxy-9-methylene), which enhance chemical stability and/or selectivity of action; and omega-chain modifications (e.g., 18,19,20-trinor-17-phenyl, 17,18,19,20-tetranor-16-phenoxy), which enhance selectivity of action and reduce biological metabolism. Derivatives of these prostaglandins include all pharmaceutically acceptable salts and esters, which may be attached to the 1-carboxyl group or any of the hydroxyl groups of the prostaglandin by use of the corresponding alcohol or organic acid reagent, as appropriate. It should be understood that the terms "analogues" and "derivatives" include compounds which exhibit functional and physical responses similar to those of prostaglandins per se.

The following publications disclose examples of prostaglandins which are suitable for use in the present invention: Crabbe, P. (ed), "Prostaglandin Research," Academic Press, New York: 1977; Advances in Prostaglandin, Thromboxane, and Leukotriene Research, 14: 263-307 (1985); ibid., 14: 309-425 (1985); US 3,884,969 (Schaub et al.); US 3,873,607 (Bernady et al.); GB 1 444 971 (Floyd, Jr. et al.); US 4,110,368 (Floyd, Jr. et al.); US 4,291,175 (Wissner et al.); US 4,321,405 (Weiss); US 4,343,949 (Bernady et al.); US 4,614,825 (Snitman et al.); US 4,029,681 (Smith); US 4,097,489 (Bundy); US 4,288,616 (Sih); US 3,755,426 (Strike et al.); US 4,576,962 (Matthews); US 4,599,353 (Bito); EP 364 417 (Stjernschantz et al.); DE 39 23 797 (Klar et al.); WO 85/02841 (Skuballa et al.); EP 299 914 (Buchmann et al.); EP 399 839 (Woodward et al.); US 4,994,274 (Chan et al.); WO 91/14428 (Woodward); US 5,093,329 (Woodward); EP 289 349 (Ueno et al.); and EP 366 279 (Ueno et al.). All of these publications are incorporated by reference herein with respect

-10-

to their disclosures and teachings concerning prostaglandin structure, synthesis and activity. It is to be understood that the prostaglandins disclosed in and taught by the above-referenced publications are only exemplary in nature; the present invention is not intended to be limited by the disclosures and teachings of the above-referenced publications.

Specific examples of prostaglandins which are useful in the present invention include: PGF_{2α}, PGE₂, PGE₁, prostacyclin, 15(S)-methyl-PGF_{2α}, 16,16-dimethyl-PGF_{2α}, 15(S)-methyl-PGE₂, 16,16-dimethyl-PGE₂, 17,18,19,20-tetranor-16-phenoxy-PGE₂, 17,18,19,20-tetranor-16-phenoxy-PGF_{2α}, 18,19,20-trinor-17-phenyl-PGE₂, 18,19,20-trinor-17-phenyl-PGF_{2α}, trimoprostil, RS-84-135, rioprostil, S-1033 (15-deshydroxy PGF_{2α}, sodium salt), S-747260, nocioprost, CS-412, YPG-209, K-10134, cloprostenol, fluprostenol, luprostiol, etiproston, tiaprost, SQ 27986, ZK 138519, 13,14-dihydro ZK 138519, ZK 118182, 13,14-dihydro ZK 118182, ZK 110841, 13,14-dihydro ZK 110841, PhXA41 (latanoprost), RO-221327, HR-466, HR-601, ONO-1206, UFO-21, 11-deoxy-PGE₂, 11-deoxy-PGF_{2α}, 11-deoxy-16,16-dimethyl-PGE₂, 11-deoxy-15(S)-methyl-PGE₂, 11-deoxy-15(S)-methyl-PGF_{2α}, misoprostol, enisoprost, MDL-646, CL-115,574, CL-115,347, TR-4161, TR-4752, TR-4367, CP-27987, sulprostone, gemeprost, alfadostrol, delprostene, prostalene, fenprostalene, CL-116,069, ONO-995, RO-229648, as well as their pharmaceutically acceptable esters and salts, as appropriate for the respective individual structures. The most preferred prostaglandins are: PGF_{2α}-1-isopropyl ester, PGF_{2α}-1-ethyl ester, RO-229648, SQ 27986, ZK 138519, 13,14-dihydro ZK 138519, ZK 110841, 13,14-dihydro ZK 110841, PhXA41 and 18,19,20-trinor-17-phenyl-PGF_{2α}-1-methyl ester. All of the foregoing compounds are known.

In general, compositions of the present invention will include one or more clonidine derivatives in an amount between about 0.02 and about 2.0 percent by weight (wt%) and one or more prostaglandins in an amount between about 0.00001 and about 0.2 wt%. It is preferred to use one or more clonidine derivatives in an amount between about 0.05 and about 1.0 wt%, and it is especially preferred to use an amount between about 0.1 and about 0.25 wt%. It is preferred to use one

-11-

or more prostaglandins in an amount between about 0.0001 and about 0.01 wt%, depending on the potency of the prostaglandin. The ratio by weight of clonidine derivative to prostaglandin is generally between about 1:1 to about 10,000:1 and preferably between about 5:1 to about 1000:1. It should be understood that the ratio by weight of clonidine derivative to prostaglandin will greatly depend on the potency of the prostaglandin used, since the potency of different prostaglandins may differ by as much as a factor of 10⁵.

In addition to the above-described principal active ingredients, the antiglaucoma compositions of the present invention may further comprise various formulatory ingredients, such as antimicrobial preservatives and tonicity agents. Examples of suitable antimicrobial preservatives include: benzalkonium chloride, thimerosal, chlorobutanol, methyl paraben, propyl paraben, phenylethyl alcohol, edetate disodium, sorbic acid, Polyquad®, Dymed® and other agents equally well known to those skilled in the art. Such preservatives, if utilized, will typically be employed in an amount between about 0.001 and about 1.0 wt%. Examples of suitable agents which may be utilized to adjust the tonicity or osmolality of the formulations include sodium chloride, potassium chloride, mannitol, dextrose, glycerine and propylene glycol. Such agents, if utilized, will be employed in an amount between about 0.1 and about 10.0 wt%.

The compositions of the present invention may additionally include components to provide sustained release and/or comfort. Such components include high molecular weight, anionic mucomimetic polymers, gelling polysaccharides and finely-divided drug carrier substrates. These components are discussed in greater detail in US 4,911,920 (Jani et al.) and in US 5,212,162 (Mssel et al.). The entire contents of these two patents are incorporated herein by reference.

The high molecular weight, anionic mucomimetic polymers useful in the present invention have a molecular weight between about 50,000 and 6 million daltons. The polymers are characterized as having carboxylic acid functional

groups and preferably contain between 2 and 7 carbon atoms per functional group. The gels which form during preparation of the ophthalmic polymer dispersion have a viscosity between about 1,000 to about 300,000 centipoise (cps). Suitable polymers are carboxy vinyl polymers, preferably those called Carbomers, e.g., 5 Carbopol® (B.F. Goodrich Co., Cleveland, Ohio). Specifically preferred are Carbopol® 934 and 940. Such polymers will typically be employed in an amount between about 0.05 and about 8.0 wt%, depending on the desired viscosity of the composition. Pourable liquid compositions generally comprise an amount of the polymer between about 0.05 and about 2.0 wt%.

10 As used herein, the term "finely-divided drug carrier substrate" (or "DCS") means finely-divided solids, colloidal particles, or soluble polymers and/or polyelectrolytes which are capable of selective adsorption or binding with drug molecules. Examples of DCS include, but are not limited to: finely divided silica, such as fumed silica, silicates and bentonites; ion exchange resins, which can be 15 anionic, cationic or non-ionic in nature; and soluble polymers, such as, alginic acid, pectin, soluble carrageenans, Carbopol®, and polystyrene sulfonic acid. In general, the DCS component is used at a level in the range of about 0.05 to about 10.0 wt%. For particulate DCS, the average particle size diameter ranges from 1 to 20 microns. The amount of DCS and its characteristics (e.g., amount of cross-linking, particle size) may be varied in order to produce the desired time-release 20 profile for the chosen drug.

Preferred DCS are the ion exchange resins. Some resins which are used in chromatography make ideal DCS for binding drugs in the compositions of the present invention. Such resins are readily available, for example, from Rohm & 25 Haas (Philadelphia, Pennsylvania) under the name Amberlite® and from Dow Chemical Co. (Midland, Michigan) under the name Dowex®. The average particle size of the commercially available forms of the resins is about 40 to 150 microns. As the particle size of the resin is critical, such commercially available particles are most conveniently reduced to a particle size range of about 1.0 to 25 microns by 30 ball milling, according to known techniques. At least 95% of the resulting

spheroidal particles must have a diameter less than 20 microns. The ion exchange resins will typically be present in an amount between about 0.05 and about 10.0 wt% and will have an average particle size diameter between about 1 and about 20 microns.

5 As will be appreciated by those skilled in the art, the compositions may be formulated in various dosage forms suitable for topical ophthalmic delivery, including solutions, suspensions, emulsions, gels and erodible solid ocular inserts. The compositions are preferably aqueous, have a pH between 3.5 to 8.0 and an osmolality between 280 to 320 milliOsmoles per kilogram (mOsm/kg).

10 The compositions of the present invention may also comprise non-aqueous formulations such as: substantially non-aqueous liquids, substantially non-aqueous semi-solid compositions and solid compositions or devices. The first class, substantially non-aqueous liquids, comprise a combination of a clonidine derivative of formula (I) and at least one prostaglandin ("drug combination") dissolved or suspended in one or more of the following: vegetable and mineral oils, such as, liquid petrolatum, corn oil, castor oil, sesame oil and peanut oil; triglycerides, such as the capric/caprylic triglycerides commonly used in foods and cosmetics; liquid lanolin and lanolin derivatives; and perfluorohydrocarbons. The second class, semi-solid compositions, comprise a drug combination dissolved or suspended in 15 one or more of the following: various types of petrolatum, such as white, yellow, red and so on; lanolin and lanolin derivatives; gelled mineral oil having a hydrocarbon base, such as Plastibase®; petrolatum and ethylene carbonate mixtures; petrolatum in combination with surfactants and polyglycol, such as 20 polyoxyl 40 stearate and polyethylene glycol.

25 The third class, solid compositions or devices, include non-erodible devices which are inserted into the conjunctival sac of the eye and later removed, such as the Alza-type diffusion or osmotic pressure controlled polymer membranes; and bioerodible polymers which do not have to be removed from the conjunctival sac, such as essentially anhydrous but water soluble polymers and resins (e.g.,

-14-

- celluloses, polycarboxylic acids, and so on). Especially preferred are the bioerodible inserts described and detailed in US 4,540,408 (Lloyd) and US 4,730,013 (Bondi et al.), wherein drug combinations of the present invention would be entrained in a non-aqueous matrix consisting essentially of polyvinyl alcohol.
- 5 The entire contents of these two patents are incorporated herein by reference.

The present invention is also directed to methods of treating glaucoma and other ophthalmic diseases and abnormalities. The methods comprise topically applying to the affected eye(s) of the patient a therapeutically effective amount of a composition according to the present invention. The frequency and amount of dosage will be determined by the clinician based on various clinical factors. The methods will typically comprise topical application of one or two drops (approximately 30 microliters) of a liquid composition, or an equivalent amount of a solid or semi-solid dosage form, to the affected eye one to two times per day.

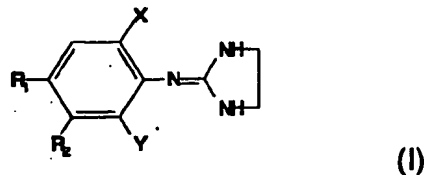
10 The invention has been described by reference to certain preferred embodiments; however, it should be understood that it may be embodied in other specific forms or variations thereof without departing from its spirit or essential characteristics. The embodiments described above are therefore considered to be illustrative in all respects and not restrictive, the scope of the invention being indicated by the appended claims rather than by the foregoing description.

What is Claimed is:

1. A topical ophthalmic composition for the treatment of glaucoma, comprising a combination of a pharmaceutically effective amount of a prostaglandin and a pharmaceutically effective amount of a clonidine derivative.

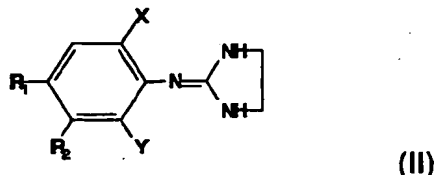
5 2. The composition of claim 1, wherein the clonidine derivative is selected from the group consisting of:

a) a compound of formula:



10 wherein: R_1 and R_2 are selected from H, OH, NHR' and O-C(=O)-CH₂-R', with R' being selected from H and C₁-C₄ alkyl, provided that one of R_1 and R_2 is hydrogen; and X and Y are selected from Br, Cl, CH₃ and CH₂CH₃; and

b) a compound of formula:



15 wherein: X and Y are selected from Br, Cl, CH₃ and CH₂CH₃, with the provision that at least one of X and Y is alkyl.

20 3. The composition of claim 1, wherein the final composition concentration of clonidine derivative is between about 0.02 and about 2.0 wt% and the final composition concentration of prostaglandin is between about 0.00001 and about 0.2 wt%.

-16-

4. The composition of claim 3, wherein the final composition concentration of clonidine derivative is between about 0.05 and about 1.0 wt%.

5. The composition of claim 4, wherein the final composition concentration of clonidine derivative is between about 0.1 and about 0.25 wt%.

5 6. The composition of claim 3, wherein the final composition concentration of prostaglandin is between about 0.0001 and about 0.01 wt%.

7. The composition of claim 1, wherein the prostaglandin is selected from the group consisting of: PGF_{2α}, PGE₂, PGE₁, prostacyclin, 15(S)-methyl-PGF_{2α}, 16,16-dimethyl-PGF_{2α}, 15(S)-methyl-PGE₂, 16,16-dimethyl-PGE₂,
10 17,18,19,20-tetranor-16-phenoxy-PGE₂, 17,18,19,20-tetranor-16-phenoxy-PGF_{2α}, 18,19,20-trinor-17-phenyl-PGE₂, 18,19,20-trinor-17-phenyl-PGF_{2α}, trimoprostil, RS-84-135, rioprostil, S-1033, S-747260, noclaprost, CS-412, YPG-209, K-10134, cloprostenol, fluprostenol, luprostiol, etiproston, tiaprost, SQ 27986, ZK 138519, 13,14-dihydro ZK138519, ZK 118182, 13,14-dihydro ZK 118182, ZK 110841, 13,14-dihydro ZK110841, PhXA41, RO-221327, HR-466, HR-601, ONO-1206, UFO-21,
15 11-deoxy-PGE₂, 11-deoxy-PGF_{2α}, 11-deoxy-16,16-dimethyl-PGE₂, 11-deoxy-15(S)-methyl-PGE₂, 11-deoxy-15(S)-methyl-PGF_{2α}, misoprostol, enisoprost, MDL-646, CL-115,574, CL-115,347, TR-4161, TR-4752, TR-4367, CP-27987, sulprostone, gemeprost, alfadprostol, delprostene, prostalene, fenprostalene, CL-116,069,
20 ONO-995, RO-229648, and their pharmaceutically acceptable esters and salts, as appropriate.

8. The composition of claim 7, wherein the prostaglandin is selected from the group consisting of: PGF_{2α}-1-isopropyl ester, PGF_{2α}-1-ethyl ester, RO-229648, SQ 27986, ZK 138519, 13,14-dihydro ZK138519, ZK 110841, 13,14-dihydro ZK 110841, PhXA41 and 18,19,20-trinor-17-phenyl-PGF_{2α}-1-methyl ester.

25 9. The composition of any of claims 1-8, further comprising an anionic, mucomimetic polymer and a finely divided drug carrier substrate.

-17-

- 10. The composition of any of claims 1-8, further comprising a gelling polysaccharide and a finely divided drug carrier substrate.**

- 11. Use of a composition of any of claims 1-10 to treat glaucoma.**

INTERNATIONAL SEARCH REPORT

Int'l. Appl. No.
PCT/US 93/09742

A. CLASSIFICATION OF SUBJECT MATTER
IPC 5 A61K31/557 // (A61K31/557, 31:415)

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
IPC 5 A61K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	EP,A,0 458 589 (KABUSHIKI KAISHA UENO SEIYAKU OYO KENKYUJO) 27 November 1991 see page 7, line 5 - line 17 ---	1-11
A	FORTSCHRITTE DER OPHTHALMOLOGIE vol. 87, no. SUPP , 1990 , HEIDELBERG pages S172 - S174 KRIEGELSTEIN G.K. 'Medikamentöse Glaukomtherapie' see page S174, column 1, line 26 - column 2, line 22 ---	1-11
A	EP,A,0 286 903 (THE TRUSTEES OF COLUMBIA UNIVERSITY IN THE CITY OF NEW YORK) 19 October 1988 see column 2, line 14 - column 3, line 34 -----	1-11

Further documents are listed in the continuation of box C.

Patent family members are listed in annex.

* Special categories of cited documents :

- 'A' document defining the general state of the art which is not considered to be of particular relevance
- 'E' earlier document but published on or after the international filing date
- 'L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- 'O' document referring to an oral disclosure, use, exhibition or other means
- 'P' document published prior to the international filing date but later than the priority date claimed

'T' later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

'X' document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

'Y' document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.

'&' document member of the same patent family

1

Date of the actual completion of the international search

Date of mailing of the international search report

17 January 1994

25.01.94

Name and mailing address of the ISA

European Patent Office, P.B. 5818 Patentiaan 2
NL - 2280 HV Rijswijk
Tel. (+ 31-70) 340-2040, Tx. 31 651 epo nl,
Fax (+ 31-70) 340-3016

Authorized officer

LEHERTE, C

INTERNATIONAL SEARCH REPORT

Information on patent family members

Int. Search Application No

PCT/US 93/09742

Patent document cited in search report	Publication date	Patent family member(s)		Publication date
EP-A-0458589	27-11-91	CA-A-	2042934	23-11-91
		JP-A-	4253912	09-09-92
EP-A-0286903	19-10-88	AU-B-	607981	21-03-91
		AU-A-	1387088	06-10-88
		DE-A-	3871596	09-07-92
		JP-A-	63313728	21-12-88
		US-A-	4952581	28-08-90

THIS PAGE BLANK (USPTO)